Distinct response of yeast ribosomes to a miscoding event during translation

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ABSTRACT
Numerous mechanisms have evolved to control the accuracy of translation, including a recently discovered retrospective quality control mechanism in bacteria. This quality control mechanism is sensitive to perturbations in the codon:anticodon interaction in the P site of the ribosome that trigger a dramatic loss of fidelity in subsequent tRNA and release factor selection events in the A site. These events ultimately lead to premature termination of translation in response to an initial miscoding error. In this work, we extend our investigations of this mechanism to an in vitro reconstituted Saccharomyces cerevisiae translation system. We report that yeast ribosomes do not respond to mismatches in the P site by loss of fidelity in subsequent substrate recognition events. We conclude that retrospective editing, as initially characterized in Escherichia coli, does not occur in S. cerevisiae. These results highlight potential mechanistic differences in the functional core of highly conserved ribosomes.

Keywords: eukaryotic ribosome; translation; fidelity; miscoding; termination

INTRODUCTION
The information contained in an organism’s genome must be faithfully translated to produce functional proteins, which in turn allows the organism to grow and reproduce. Perfect fidelity, however, has a price: It would be too slow, and too costly in energetic terms, for a living organism (Thompson and Karim 1982; Ruusala et al. 1984). Hence, all branches of life have arrived at some compromise between speed and accuracy in transmission that allows for optimal survival.

This evolved level of fidelity during translation is not the same in all organisms; indeed, more complex organisms appear to have greater overall fidelity than simpler ones. For example, the in vivo miscoding frequencies in yeast appear to be between $1 \times 10^{-4}$ and $5 \times 10^{-4}$ misreading events per codon (Stansfield et al. 1998; Rakwalska and Rospert 2004; Salas-Marco and Bedwell 2005; Plant et al. 2007; Kramer et al. 2010), about 10-fold less frequent than the miscoding frequencies measured in vivo in Escherichia coli (Parker 1989; Kramer and Farabaugh 2007). In higher eukaryotes errors may be even less frequent (Martin et al. 1989).

These numbers provide a sense for the eventual outcome from whatever mechanisms exist to ensure this fidelity, but do not define the molecular process. For some time, much biochemical effort has been focused on understanding how the ribosome carefully selects the appropriate aminoacyl-tRNA (or release factor) during each round of elongation (or termination) (for review, see Zaher and Green 2009a). And, while this selection process appears to account for a majority of the discrimination at the level of elongation and termination in bacteria, there also appear to be molecular mechanisms in place, at least in certain bacteria, to retrospectively (after peptide bond formation) monitor (and increase) fidelity (Zaher and Green 2009b). This quality control mechanism monitors codon:anticodon interactions in the P and E sites of the ribosome, where the presence of mismatches appears to increase the promiscuity of interactions in the A site of the ribosome. The loss of A site fidelity manifests itself in one of two ways: Either another incorrect tRNA is selected, thereby iterating the miscoding error, or a release factor is selected at a sense codon, leading to premature termination. The ultimate consequence of this process is that protein synthesis is prematurely terminated in response to one or more miscoding errors (Zaher and Green 2009b). This retrospective quality control system...
is somewhat reminiscent of the proofreading activity of DNA polymerase in that both increase the fidelity of the process, though in the latter case the mistakes are actually repaired.

Here we explored the fidelity of codon selection by both tRNAs and release factors in our newly developed in vitro reconstituted yeast translation system. We focused our analysis on decoding events that occur on ribosome complexes with fully matched as well as mismatched P-site codon:anticodon helices. In broad terms, tRNA selection appears to share many similarities with the comparable events in *E. coli*, including, for example, increased miscoding and decreased release activity in the presence of aminoglycosides. Surprisingly, however, we saw only modest responses of the yeast ribosome to mismatches in the P site, and these responses would not obviously increase the overall fidelity of translation. Consequently, we conclude that retrospective quality control as initially defined in *E. coli* does not take place in yeast. It remains possible, however, that post-PT quality control does occur in eukaryotes, through a distinct mechanism, potentially involving extra-ribosomal factors.

**RESULTS**

**Experimental system**

For this study, we developed an in vitro reconstituted translation system with *Saccharomyces cerevisiae* components that could specifically follow the events of translation elongation and termination. Initiation complexes were prepared from purified yeast subunits (40S and 60S) and short unstructured mRNAs with a core set of recombinant initiation factors (eIF1, eIF1A, eIF5, eIF5B, eIF2, and eEF1A) charged in vitro transcribed initiator methionine tRNA) as previously described (Acker et al. 2007).

For elongation reactions, the translation factors eEF1A, eEF2, and eEF3 were expressed and purified as previously described (Jorgensen et al. 2002; Andersen et al. 2004). Commercially available tRNA<sup>Phe</sup> and in vitro transcribed tRNA<sup>Lys</sup> (see Materials and Methods) were chosen as elongation substrates, as both were efficiently aminoacylated by an S100 extract and incorporated by the ribosome when provided in a ternary complex with eEF1A•GTP.

With our first set of experiments, we tested the behavior of the system during elongation. Initiation complexes were prepared on short mRNAs containing the coding sequence AUG UUC AAA, as described above, and the elongation cycle was performed. In the absence of ternary complex, only Met is observed (Fig. 1, lane 1). When Phe-tRNA<sup>Phe</sup>•eEF1A•GTP ternary complex is added, Met-Phe dipeptide is formed (Fig. 1, lane 2). When Lys-tRNA<sup>Lys</sup>•eEF1A•GTP ternary complex is included, synthesis of the tripeptide Met-Phe-Lys is only observed when both elongation factors eEF2 and eEF3 are also added (Fig. 1, lanes 3,4). The results from this staged reaction sequence indicate that our translation elongation system is appropriately responsive to the various components.

We next prepared reagents that would allow us to study translation termination in the yeast reconstituted system. N-terminally His-tagged versions of the class 1, eRF1, and class 2, eRF3, release/termination factors from *S. cerevisiae* were cloned, expressed in, and purified from *E. coli*. An N-terminally truncated version of eRF3 (ΔN165) (lacking a poorly behaved glutamine-rich domain) was utilized to increase solubility, as has been described in a reconstituted mammalian system (Alkalaeva et al. 2006). The activity of the termination factors was evaluated on ribosome complexes programmed with an mRNA carrying the coding sequence AUG UUC AAA UAA, Met-Phe-Lys-tRNA<sup>Lys</sup> poised in the P site, and the stop codon (UAA) in the A site. The purified release factors eRF1 and eRF3 together stimulated peptide release in the presence of GTP, while the non-hydrolyzable analog GMPPNP inhibited the release reaction (Fig. 2A). Additionally, eRF1 was able to stimulate peptide release alone, and the rate was substantially increased by the addition of eRF3 (Fig. 2B). These data are overall markedly similar to earlier observations in the reconstituted mammalian system (Alkalaeva et al. 2006).

With these reagents in place, the reconstituted yeast system allows for detailed in vitro analysis of translation elongation and termination.

**Effects of paromomycin on miscoding**

The aminoglycoside antibiotic paromomycin is known to promote translational miscoding in bacteria as a result of binding to a site in the highly conserved decoding center 16S rRNA (Moazed and Noller 1987; Carter et al. 2000). Similarly, paromomycin appears to stimulate miscoding in *S. cerevisiae* as determined with a series of reporter constructs.

![Figure 1](image-url)
Here, we evaluated the effects of paromomycin on tRNA and eRF selection in our in vitro reconstituted system. Initiation complexes were assembled on mRNAs with the terminological, there are four possible complexes that were (‘‘cognate’’ or ‘‘near cognate,’’ respectively). Given this, and with a cognate or near-cognate codon in the A site (‘‘matched’’ or ‘‘mismatched,’’ respectively) the P site (‘‘matched’’ or ‘‘mismatched,’’ respectively) are encoded on UUC encoding mRNAs, while the mismatched complexes are formed on UUG encoding mRNAs. In each case, either AAA or AAU is poised in the A site for cognate or near-cognate decoding by tRNA Lys, respectively. We confirmed the positioning of the mRNAs relative to the ribosome by toeprinting (Hartz et al. 1988) to rule out the possibility that mismatched complexes are prone to frameshifting (data not shown).

First, in a simple experiment where modest amounts of eRF1•GTP•Lys-tRNA Lys ternary complexes were added, the cognate reactions went to completion, while little mis-coding was observed with the near cognates, on both matched and mismatched complexes (Fig. 3B). We next asked whether there might be differences in the rates of these different reactions that might reflect changes in ribosome structure resulting from the perturbation of the P site. The cognate reactions were evaluated with reactions containing subsaturating amounts of Lys-tRNA Lys ternary complex (so called $k_{cat}/K_m$ conditions) and were seen to be relatively similar to one another, with $k_{obs}$ values of 0.16 min$^{-1}$ and 0.59 min$^{-1}$ on matched and mismatched complexes, respectively (Fig. 3C). The slower near-cognate reactions were evaluated using saturating amount of Lys-tRNA Lys ternary complex (so called $k_{cat}$ conditions) and were again found to be relatively similar to one another, 0.24 min$^{-1}$ and 0.65 min$^{-1}$, on matched and mismatched complexes (Fig. 3D). Most significantly, the modest increases in rate seen on the mismatched complexes were about the same for the cognate and near-cognate reactions (3.7- and 2.7-fold, respectively).

Since the effects of the P-site mismatches on cognate reactions were performed at subsaturating concentrations of ternary complex, the observed effects may represent a combination of binding and catalytic defects. By contrast, since the effects of the P-site mismatch on near-cognate reactions were performed under saturating substrate concentrations, no binding component is included in the observed data. We note that these experiments were performed differently because the fast rates of tripeptide synthesis for cognate complexes require use of a quench-flow apparatus, and we are not yet able to generate the requisite quantities of material. We think it unlikely that the results would be qualitatively different if all measurements were made under $k_{cat}$ conditions. Our results with the release factors (below) suggest that binding to the A site is generally unperturbed in the mismatched complex. Together, these data suggest that while P-site mismatches do (modestly) accelerate tRNA selection in the A site for both cognate and near-cognate tRNAs, these effects are not specific for near-cognate tRNAs, and thus do not change the overall fidelity of the selection step.

To further test this hypothesis, we conducted tRNA competition experiments in which matched and mismatched
dipeptidyl-tRNA complexes (with Met-Phe-tRNA\textsuperscript{Phe} on AUGUUCAAA and AUGUUGAAA mRNAs, respectively) were reacted with total aminoacylated tRNA from a translating extract (Wu et al. 2007). The diverse reaction products were resolved on a two-dimensional TLC system where the first phase was a passive TLC separation using 70:20:10 EtOH:H\textsubscript{2}O:HOAc as the mobile phase, and the second phase was electrophoretic using the pyridine:acetate mobile phase described earlier (Zaher and Green 2009b). Overall, the patterns of product formation are quite similar for the matched and mismatched complexes (Supplementary Figure 1); spots for Met, Met-Phe, and Met-Phe-Lys can be identified in each, in addition to several unidentified bands produced in approximately equal quantities by the distinct complexes. Consistent with our kinetic results, this competition experiment argues that mismatches in the P site do not promote iterated errors in tRNA selection in S. cerevisiae, as previously documented in E. coli (Zaher and Green 2009b).

**FIGURE 3.** Miscoding by yeast ribosomes during elongation is stimulated by paromomycin but not by P-site mismatches. (A) Met-Lys dipeptide synthesized on Met-Lys (dark bars) or Met-Asn (light bars) mRNA in the presence of increasing concentrations of paromomycin (indicated on the x-axis). (B) The extent of misreading (fraction dipeptide converted into tripeptide) is not substantially affected by P-site mismatches at moderate Lys-tRNA\textsuperscript{Lys}\textsuperscript{+} ternary complex concentrations. Complexes were prepared with Met-Phe dipeptidyl-tRNA in the P site with either a matched Phe (UUC) (dark bars) or mismatched Leu (UUG) (light bars) codon in the P site with either a cognate Lys (AAA) or a near-cognate Asn (AAU) codon in the A site (x-axis). Rates of tripeptide synthesis by (C) cognate complexes at subsaturating Lys-tRNA\textsuperscript{Lys}\textsuperscript{+} ternary complex concentration, and by (D) near-cognate complexes at saturating Lys-tRNA\textsuperscript{Lys}\textsuperscript{+} ternary complex concentration. All data are presented as mean ± standard error of at least two measurements.

**DISCUSSION**

Our quantitative analysis of the events of decoding and termination in yeast both on normal (matched) and P-site mismatched elongation and termination complexes has measuring the rates of peptide release on matched and mismatched dipeptidyl-tRNA ribosome complexes (Met-Phe-tRNA\textsuperscript{Phe}) containing a UAA stop codon in the A site (AUGUUCAAA and AUGUUGAAA coding sequence, respectively) and observed only small differences in the rates of catalysis in the presence of eRF1 alone or eRF1•eRF3 (indeed, the modest differences were in the opposite direction than anticipated) (Fig. 4A). We also directly measured the binding affinity between eRF1•eRF3 and the matched and mismatched ribosome complexes using a fluorescence assay where the P-site tRNA\textsuperscript{Phe} is modified with a proflavin residue (Wintermeyer and Zachau 1979) and intrinsic fluorescence changes are followed (Zaher and Green 2010); here again we failed to observe any significant differences (Fig. 4B). Finally, $K_{1/2}$ values for the canonical release reaction were determined as previously (Zaher and Green 2009b) for the matched and mismatched complexes; again we observed no differences in these values for the distinct ribosome complexes with either eRF1•eRF3 or eRF1 alone (Fig. 4C). These data are strikingly different than those reported in E. coli where even on stop codon containing complexes, $K_{1/2}$ differences between matched and mismatched complexes were easily discerned (Zaher and Green 2009b).

We next measured the rate constants for peptide release on matched and mismatched complexes carrying a codon in the A site that is near-cognate to a stop codon, specifically the UAC “sense” tyrosine codon. As anticipated, the rates of release observed in both reactions were exceedingly slow, with half-lives on the order of 14 h (Fig. 4D). Indeed, the observed rates of peptide release in this system are not faster than the uncatalyzed rate of the reaction (with an inactive eRF1 variant [AGQ] or in the absence of release factors) where the loss of stability in the mismatched complexes contributes to slightly elevated rates of peptidyl-tRNA dissociation and solution hydrolysis. Again, these results provide no evidence to support a model where premature termination is triggered by mismatches in the P site, as previously documented in E. coli (Zaher and Green 2009b).
revealed key similarities and differences between the molecular events of translation in bacteria and eukaryotes. To probe the biochemical features of eukaryotic translation elongation and termination, we developed an in vitro reconstituted translation system where various stalled ribosome complexes could be prepared and evaluated using presteady state kinetic approaches. As anticipated based on the known high level of translational fidelity in vivo ($\sim 10^{-12}$) (Salas-Marco and Bedwell 2005; Kramer et al. 2010), the acceptance of aminoacyl-tRNAs and eRF1 on near-cognate codons was minimal in the in vitro system. Moreover, as in bacteria, the aminoglycoside class of antibiotics stimulated misreading during tRNA selection and inhibited release factor function. These observations are consistent with the fact that the functional core of the ribosome is well conserved from bacteria to eukaryotes (Alksne et al. 1993; Gutell et al. 1994; Liebman et al. 1995).

Despite these similarities, we observed substantial differences in the response of the eukaryotic ribosome to mismatches in the P-site codon:anticodon helix relative to the dramatic response of \textit{E. coli} ribosomes (Zaher and Green 2009b). While there were modest increases in the overall rates of peptidyl transfer on ribosome complexes carrying a mismatch in the P site (suggestive of communication between the P and A site), these effects were uniform for both cognate and near-cognate decoding events, thus not impacting the overall fidelity of the subsequent round of elongation. Secondly, the rates of peptide release, and the $K_{1/2}$ and the $K_d$ of release factors for stop and near-stop codon-programmed ribosome complexes, were unaltered in the presence of the P-site mismatch. Indeed, premature termination on a sense codon appears to be a particularly unlikely outcome in yeast, given the exceedingly slow rates that we observed for this reaction ($6.5 \times 10^{-14}$ min$^{-1}$; Fig. 4D). Given these two different sets of observations, we argue that retrospective quality control (as mediated through iterated miscoding and premature termination following an initial misreading event) does not happen in yeast as it does in \textit{E. coli}.

We are cautious in emphasizing that these results do not exclude the possibility that some form of retrospective monitoring of ribosome complex integrity does happen in yeast. For example, it is possible that our experiments lack a key factor that is present in vivo and is responsible for recognition of the P-site mismatched ribosome complexes, though we note that we did not observe any premature termination of matched or mismatched complexes when cellular extracts were added (data not shown).

It is noteworthy that eukaryotes exhibit greater overall translational fidelity, apparently without retrospective quality control that bacteria exhibit with it. Hence, our observations lead us to speculate about what might be the reason for such fundamental differences in quality control during translation in bacteria and eukaryotes. On the one hand, the phenomena that we have characterized in bacteria appears to simply report on structural perturbations in the A site of the ribosome that result from triggering structural perturbations in the neighboring P (and E) site(s). These A-site perturbations are manifested as promiscuous tRNA and RF selection activity that ultimately leads to premature termination of protein synthesis. While the premature termination event in bacteria is unusually sensitive to paromomycin by a pair of mutations in the decoding site and peptidyl transferase center are highly conserved, and their mutation is lethal in organisms in both kingdoms (Cochella et al. 2007; Fan-Minogue and Bedwell 2008). Still, yeast and bacterial ribosomes are not identical even in these functionally critical regions, and some of the molecular differences have significant consequences for ribosome function. For example, paromomycin resistance in yeast is mediated by a pair of nucleotides in helix 44 in the decoding site; yeast can be rendered sensitive to paromomycin by a pair of mutations.
A1754G and G1645A that restore the bacterial sequence to helix 44 (Fan-Minogue and Bedwell 2008). It seems possible that features of the ribosome that are important for retrotranscription editing may be missing from yeast ribosomes, either because these features arose late in evolution in bacteria or were lost from yeast after these branches of the tree diverged.

It is also possible that retrotranscription quality control in bacteria is promoted by special features of both the ternary complex and the release factors in recognizing mismatch-carrying ribosome complexes. Bacterial and eukaryotic release factors are completely different, independently evolved proteins and, as such, it is possible that they might respond differently to mismatches in the P site. In contrast, EFTu and eEF1A are homologous proteins that function similarly in tRNA selection in the two systems (for example, paromomycin promotes miscoding in both systems), making this factor unlikely to be the basis for the observed differences. Moreover, since commercially available yeast tRNA<sup>Phe</sup> was used for many of the experiments in both the bacterial (Zaher and Green 2009b, 2010) and the yeast experiments, tRNAs per se are unlikely to explain the different responses seen in bacteria and eukaryotes.

Another possible explanation for the lack of retrotranscription editing in yeast is that fidelity is more effectively imposed during the kinetically slower tRNA selection process (Mathews et al. 2007). Alternatively, since retrotranscription editing may be principally involved in the control of frameshifting in bacteria, it may be that spontaneous frameshifting is a sufficiently rare event in yeast as to require no additional control mechanisms. Given that the ORFs in yeast are generally longer than in <i>E. coli</i>, the yeast may have generally evolved higher overall levels of tRNA selection fidelity to ensure the production of accurate, full-length proteins.

In conclusion, we have shown that mismatched yeast ribosome complexes behave quite distinctly from their bacterial counterparts in vitro. The consequence of the different behaviors is that yeast do not exhibit the retrotranscription quality control mechanism as it exists in bacteria. That said, we certainly wonder whether mismatched ribosome complexes are targeted by non-core translation factors, as is the case in no-go decay (Shoemaker et al. 2010) and other mRNA decay pathways. Further work in extracts and in vivo may reveal the answers to these questions.

**MATERIALS AND METHODS**

**Ribosome preparation**

A small culture of the yeast strain YAS2488 (MATa leu2-3 112 his4-539 trp1 ura3-52 cup1::LEU2/PKG1pG/MFA2pG) (Acker et al. 2007) was grown in YPD medium for 24 h at 30°C on a roller wheel. The starter culture was then diluted into 18 L of YPD, and the large cultures were grown at 30°C with shaking until they reached an OD<sub>600</sub> of 1.0. The cells were then collected by centrifugation, washed, and resuspended in a 1/10th volume of lysis buffer (1× ribo buffer A, 1 mg/mL heparin, 2 mM DTT, 7.5 mM Mg(OAc)<sub>2</sub>, 400 mM KCl, with Roche EDTA-free Complete Protease Inhibitors). Ribo buffer A was prepared as a 10× stock of 1 M KOAc, 200 mM HEPESS-KOH pH 7.4, and 25 mM magnesium acetate. The cell paste was frozen as droplets in liquid nitrogen and lysed by grinding under cryogenic conditions, using the model 6870 freezer/mill (SPEX SamplePrep). The frozen powder was stored at −80°C until use.

The lysate was thawed and clarified by centrifugation at 25,000 × g for 30 min at 4°C. The supernatant was layered on top of a sucrose cushion (1× ribo buffer A, 500 mM KCl, 1 M sucrose, 2 mM DTT, 7.5 mM Mg(OAc)<sub>2</sub>). The ribosomes were pelleted by centrifugation at 264,900 × g for 106 min at 4°C. The ribosomes were resuspended in subunit separation buffer (50 mM HEPESS-KOH pH 7.4, 2 mM MgCl<sub>2</sub>, 500 mM KCl, 2 mM DTT), stirred gently on ice for ~30 min, then clarified by a 1-min spin at maximum speed in a refrigerated microfuge.

Subunits were separated by treatment with 1 mM puromycin. The ribosome solution was layered onto a 5%–20% sucrose gradient and centrifuged in a Beckman SW28 rotor at 76,221 × g for 9 h at 4°C. Separated 40S and 60S subunits were collected with the use of an in-line UV detector. Subunit-containing fractions were pooled and concentrated, and the buffer was exchanged to ribosome storage buffer (1× ribo buffer A, 250 mM sucrose, 2 mM DTT). Aliquots of the purified subunits were stored at −80°C until use.

**Initiation and elongation factors**

Our system utilizes the initiation factors eIF1, eIF1A, eIF5, eIF5B, and eIF2. We purified these factors according to published methods (Acker et al. 2007). The EF-Tu homolog, eEF1A, was purified in its native form from the same strain we used as the source of our ribosomes. A post-ribosomal supernatant was prepared in buffer with 50 mM KCl (buffer: 20 mM Tris pH 7.5, 0.1 mM EDTA, 1 mM DTT, 25% glycerol), and bound to DE52 resin. The unbound fraction was bound to CM-Sepharose. The bound fraction from the CM-Sepharose was eluted with 300 mM KCl, diluted to 50 mM KCl, and loaded onto a pre-equilibrated Tricorn column (Source 15S, 4.6/100 PE, GE Healthcare). Bound proteins were eluted with a linear gradient to 300 mM KCl. The eEF1A-containing fractions were identified by SDS-PAGE, pooled, and dialyzed against buffer with 100 mM KCl overnight. His-tagged eEF2 and eEF3 proteins were purified according to published procedures (Jorgensen et al. 2002; Andersen et al. 2004), and we followed these protocols as written.

**Model mRNA**

Model mRNAs were in vitro transcribed off of DNA oligonucleotides to generate mRNAs of the form: GG(UC)<sub>8</sub>UA U GU U CA A A. A variety of different codons were placed in the second and third coding positions of the mRNA including UUC (Phe), UUG (Leu, or near-cognate Phe), AAA (Lys), AAU (Asn, or near-cognate Lys), UAA (Stop), or UAC (Tyr, or near-cognate Stop).

**Preparation and charging of tRNAs**

Phenylalanine-specific tRNA from <i>S. cerevisiae</i> was purchased either from Sigma or from Chemblock. Charging reactions contained
The coding sequence of the *S. cerevisiae* gene for eRF1 (SUP45) was amplified by PCR and cloned into pPROEX-HtB (Invitrogen). The resultant plasmid (pDE8) was transformed into *E. coli* BL21(DE3). Cultures were grown in Terrific Broth, and expression was induced with IPTG. Cells were harvested by centrifugation, washed, and stored in small aliquots at −80°C. The S100 extract was then added to the 80S initiation complex and allowed to react for 5 min. If initiation complexes are desired, they can be pelleted at this point, as will be described later.

For elongation, an eEF1A ternary complex was prepared by combining 25 pmol Phe-tRNA^Phe^, 50 pmol eEF1A, and 1 mM GTP in 15 μL, and preincubating for 15 min. This ternary complex was then added to the 80S initiation complex and allowed to react for 5 min. If Phe- or Leu-miscoding is desired, the magnesium concentration of the reaction was adjusted to 10 mM. Translocation was accomplished by adding GTP (1 mM final), ATP (1 mM final), 40 pmol eEF2, and 40 pmol eEF3, and incubating for an additional 5 min.

The stalled elongation complex thus generated was layered onto a gel filtration buffer (Buffer E with 20%, sucrose). The reactions were centrifuged at 26,000 g for 1 h at 4°C in a Beckman MA-130 rotor. The ribosome pellets were resuspended in 50 μL buffer E, and then stored in single-use aliquots at −80°C.

### Tripeptide synthesis assay

Lys-tRNA^Met^, 1 mM GTP, and 10 pmol of eEF1A were combined in 10 μL of 100 mM potassium acetate buffer, pH 7.6, 2 mM DTT, 10% glycerol. Fractions containing eRF3 were pooled, concentrated, and aliquoted at −80°C.

### Release assays

Release factor ternary complex was assembled by combining 10 pmol eRF1, 60 pmol of eRF3AN165, and 1 mM GTP in a volume of 15 μL for 5 min. An aliquot of pretermination complex was thawed and added to the RF ternary complex to start the reaction. Reaction aliquots were removed and quenched with formic acid. The quenched samples were stored on ice, and are spotted and run on electrophoretic TLCs as above.

1× buffer 517 (30 mM HEPES-KOH pH 7.4, 30 mM KCl, 15 mM MgCl₂, 4 mM ATP, 10 μM phenylalanine, 5 mM DTT, and a 1/10th volume of an S100 extract. Reactions were incubated for 15 min at 37°C, then extracted twice with acid-buffered phenol and once with chloroform. Nucleic acids were precipitated with ethanol, resuspended in 20 mM KOAc, 2 mM DTT, pH 5.2, and stored in small aliquots at −80°C. The S100 extract was prepared as follows: Yeast cells were grown and lysed essentially as for the ribosome preparation. The lysis buffer used was 10 mM potassium phosphate buffer, pH 7.2, with the Roche EDTA-free Complete Protease Inhibitor cocktail. The post-ribosomal supernatant was applied to DE52 resin, and bound proteins were eluted with 250 mM potassium phosphate, pH 6.5. The eluate was concentrated, glycerol was added to a final concentration of 5%, and DTT was added to a final concentration of 2 mM. The extract was aliquoted and stored at −80°C.

Initiator methionine tRNA was in vitro transcribed off of a plasmid and charged as described in published literature (Acker et al. 2007).

Lysine-specific tRNA was in vitro transcribed off of a plasmid prepared in our laboratory. This tRNA was charged according to the same protocol as the tRNA^Phe^ described above.

### Release factors

The powder was resuspended in lysis buffer, clarified by centrifugation at 20,000 × g, and applied to a pre-equilibrated 5 mL HisTrap FF column (GE Healthcare). Bound proteins were eluted with elution buffer (lysis buffer with 500 mM imidazole). The eluate was concentrated, glycerol was added to a final concentration of 5%, and DTT was added to a final concentration of 2 mM. The extract was aliquoted and stored at −80°C.

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### Response of yeast ribosomes to miscoding errors

We assembled our ribosome complexes and carried out all our assays in buffer E (20 mM Tris-Cl pH 7.5, 100 mM KOAc, 2.5 mM Mg(OAc)₂, 0.25 mM spermidine, and 2 mM DTT) (Alkalaeva et al. 2006). All reactions take place at 26°C unless otherwise noted. Ribosome complex assembly began with the preparation of ternary complex. For ternary complex preparation, 20 pmol of charged Met-tRNA^Met^, 1 mM GTP, and 40 pmol of eRF2 were combined in 10 μL and incubated for 15 min. In a second tube, the 43S components were assembled: 40 pmol of 40S subunits, 400 pmol of model mRNA, 200 pmol of eIF1, and 100 pmol of eIF1A, in a total volume of 10 μL. The 43S components were added to the ternary complex and incubated for 5 min to generate 43S complex. In a fresh tube, the 80S components were assembled: 50 pmol 60S subunits, 100 pmol eIF5, 80 pmol eIF5B, and 1 mM GTP, in 10 μL. The 80S components were added to the 43S complex and incubated for 1 min to generate 80S initiation complexes. If initiation complexes are desired, they can be pelleted at this point, as will be described later.

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The stalled elongation complex thus generated was layered onto a gel filtration buffer (Buffer E with 35% sucrose). The reactions were centrifuged at 263,970 × g for 1 h at 4°C in a Beckman MLA-130 rotor. The ribosome pellets were resuspended in 50 μL buffer E, and then stored in single-use aliquots at −80°C.

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Lys-tRNA^Met^, 1 mM GTP, and 10 pmol of eEF1A were combined in 10 μL of 100 mM potassium acetate buffer, pH 7.6, 2 mM DTT, 10% glycerol. Fractions containing eRF3 were pooled, concentrated, and aliquoted at −80°C.

We assembled our ribosome complexes and carried out all our assays in buffer E (20 mM Tris-Cl pH 7.5, 100 mM KOAc, 2.5 mM Mg(OAc)₂, 0.25 mM spermidine, and 2 mM DTT) (Alkalaeva et al. 2006). All reactions take place at 26°C unless otherwise noted. Ribosome complex assembly began with the preparation of ternary complex. For ternary complex preparation, 20 pmol of charged Met-tRNA^Met^, 1 mM GTP, and 40 pmol of eRF2 were combined in 10 μL and incubated for 15 min. In a second tube, the 43S components were assembled: 40 pmol of 40S subunits, 400 pmol of model mRNA, 200 pmol of eIF1, and 100 pmol of eIF1A, in a total volume of 10 μL. The 43S components were added to the ternary complex and incubated for 5 min to generate 43S complex. In a fresh tube, the 80S components were assembled: 50 pmol 60S subunits, 100 pmol eIF5, 80 pmol eIF5B, and 1 mM GTP, in 10 μL. The 80S components were added to the 43S complex and incubated for 1 min to generate 80S initiation complexes. If initiation complexes are desired, they can be pelleted at this point, as will be described later.

For elongation, an eEF1A ternary complex was prepared by combining 25 pmol Phe-tRNA^Phe^, 50 pmol eEF1A, and 1 mM GTP in 15 μL, and preincubating for 15 min. This ternary complex was then added to the 80S initiation complex and allowed to react for 5 min. If Phe- or Leu-miscoding is desired, the magnesium concentration of the reaction was adjusted to 10 mM. Translocation was accomplished by adding GTP (1 mM final), ATP (1 mM final), 40 pmol eEF2, and 40 pmol eEF3, and incubating for an additional 5 min.

The stalled elongation complex thus generated was layered onto a gel filtration buffer (Buffer E with 35% sucrose). The reactions were centrifuged at 263,970 × g for 1 h at 4°C in a Beckman MLA-130 rotor. The ribosome pellets were resuspended in 50 μL buffer E, and then stored in single-use aliquots at −80°C.

Lysine-specific tRNA was in vitro transcribed off of a plasmid and charged as described in published literature (Acker et al. 2007).

Lysine-specific tRNA was in vitro transcribed off of a plasmid prepared in our laboratory. This tRNA was charged according to the same protocol as the tRNA^Phe^ described above.
Release factor binding assays

Dipeptide pretermination complexes were assembled as described above, except that the Phe-tRNA^Phe used is labeled with proflavin at positions 16 and 17. Labeling was done according to published protocols (Wintermeyer and Zachau 1979), and the labeled tRNA was charged as described above. For the titrations, ~10 nM complex, 0.5 mM GTP, and 1 μM eRF3ΔN165 were combined in a cuvette, and the fluorescent signal from the proflavin was measured. Increasing amounts of eRF1 were added, and the fluorescence was measured. The change in fluorescence was plotted as a function of eRF1 concentration and fit to a hyperbola to calculate K_d.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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